QP5030 US NA CLAIMS What is claimed is: A process for increasing the relative viscosity (RV) of a reactant 5 polyamide, comprising: contacting the reactant polyamide and a chain extender compound selected from the group consisting of bis-N-acyl bislactam compounds and mixtures thereof, both in a liquid phase, the reactant polyamide having a starting RV, a concentration of amine end groups (AEG(R)), and a concentration of carboxyl end groups (CEG(R)) with the AEG(R) being 10 greater than the CEG(R), the chain extender compound having a concentration of lactam end groups (LEG) of at least about 10 equivalents per million grams of the reactant polyamide less than the AEG(R), for a duration of about 0.5 minutes to about 10 minutes, increasing the RV of the reactant polyamide; 15 forming a product, other than flake; and quenching the product to room temperature such that the product has a final RV greater than the starting RV and a concentration of amine end groups (AEG(P)) as defined by the formula: 20 $AEG(P) = \{AEG(R) - LEG\} \pm X$ wherein: AEG(P) is the concentration of amine end groups in the quenched product in units of equivalents/1,000,000 gms of the product; AEG(R) is the concentration of amine end groups in the reactant 25 polyamide in units of equivalents/1,000,000 gms of the reactant polyamide; LEG is the concentration of lactam end groups in the chain extender compound in units of equivalents/1,000,000 gms of the reactant polyamide; and X is any number in a range of about 0 to about 5. 30 2. The process of Claim 1, wherein X is any number in a range of about 0 to about 2. The process of Claim 1, wherein the chain extender compound is 35 3. selected from the group consisting of bis-N-acyl bis-caprolactam compounds and mixtures thereof. - 25 -

QP5030 US NA

5

10

15

20

25

30

35

- 4. The process of Claim 1, wherein the chain extender compound is selected from the group consisting of isophthaloyl bis-caprolactam (IBC), adipoyl bis-caprolactam (ABC), terphthaloyl bis-caprolactam (TBC), and mixtures thereof.
- 5. The process of Claim 1, wherein the polyamide is selected from the group consisting of poly(hexamethylene adipamide) homopolymer, poly(e-caproamide) homopolymer, polydodecanolactam homopolymer, poly(tetramethyleneadipamide) homopolymer, poly(hexamethylene sebacamide) homopolymer, the polyamide of n-dodecanedioic acid and hexamethylenediamine homopolymer, the polyamide of dodecamethylenediamine and n-dodecanedioic acid homopolymer, copolymers thereof, and mixtures thereof.
- 6. The process of Claim 1, wherein the forming step comprises spinning the polyamide to form the product which is at least one filament.
- 7. A product made by the process of:

contacting reactant polyamide and a chain extender compound selected from the group consisting of bis-N-acyl bislactam compounds and mixtures thereof, both in a liquid phase, the reactant polyamide having a starting RV, a concentration of amine end groups (AEG(R)), and a concentration of carboxyl end groups (CEG(R)) with the AEG(R) being greater than the CEG(R), the chain extender compound having a concentration of lactam end groups (LEG) of at least about 10 equivalents per million grams of the reactant polyamide less than the AEG(R), for a duration of about 0.5 minutes to about 10 minutes, increasing the RV of the reactant polyamide;

forming the product, other than flake; and quenching the product to room temperature such that the product has a final RV greater than the starting RV and a concentration of amine end groups (AEG(P)) as defined by the formula:

AEG(P) = $\{AEG(R) - LEG\} \pm X$ wherein:

AEG(P) is the concentration of amine end groups in the quenched product in units of equivalents/1,000,000 gms of the product;

QP5030 US NA AEG(R) is the concentration of amine end groups in the reactant polyamide in units of equivalents/1,000,000 gms of the reactant polyamide; LEG is the concentration of lactam end groups in the chain extender compound in units of equivalents/1,000,000 gms of the reactant polyamide; 5 and X is any number in a range of about 0 to about 5. A filament, comprising: 8. a synthetic melt spun polymer including: polyamide repeating units (R₁); 10 polyamide chain extender moieties (R2), each independently, selected from the group consisting of bis-N-acyl bislactam moieties; and terminal groups (R₃), each independently, selected from the group consisting of a hydrogen atom and a hydroxyl group; the polymer including chains, each independently, having a chemical 15 structure: $R_3-(R_1-R_2)_v-R_1-R_3$

wherein y is an integer of 1-7; and the filament has a formic acid relative viscosity of at least about 30.

- 9. The filament of Claim 8, wherein R_1 , each independently, is selected from the group consisting of
- (i) -{ $CO(CH_2)_k$ -CONH-(CH_2)_mNH}_n-, where k and m, each independently, is an integer of 1-12, and n is an integer of 10-140, and

25

35

- (ii) $-\{NH(CH_2)_x-CO\}_z$ where x is an integer of 1-12 and z is an integer of 20-280.
- 10. The filament of Claim 8, wherein R₂, each independently, is selected from the group consisting of bis-N-acyl bis-caprolactam moieties.
 - 11. The filament of Claim 8, wherein R₂, each independently, is selected from the group consisting of an isophthaloyl bis-caprolactam (IBC) moiety, an adipoyl bis-caprolactam (ABC) moiety, and a terphthaloyl bis-caprolactam (TBC) moiety.